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(54) Title: 2-SUBSTITUTED 1,25-DIHYDROXYVITAMIN D ₃ DERIVATIVES					

(57) Abstract

Vitamin D₃ analogues which include a 2-substituted alcohol or fluoride are described. The preferred alcohol substituent is exemplified by the structural formula -(CH₂)₄OH and the preferred fluoride substituents by the structural formulas -(CH₂)₄F and -(CH₂)₄F. Methods for the preparation of a vitamin D₃ analogue which includes a 2-substituted alcohol or fluoride starting with 2+4-cycloaddition of commercially available methyl 2-pyrone-3-carboxylate are also described.

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2-SUBSTITUTED 1,25-DIHYDROXYVITAMIN D3 DERIVATIVES

BACKGROUND OF THE INVENTION

The present invention relates to novel biologically active vitamin D₃ analogues which include an alcohol or fluoride substituent in the 2-position and methods for their preparation.

FIELD OF THE INVENTION

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Vitamin D₃ analogues have been recognized as having important biological activities. It is known, for example, that vitamin D₃ analogues can be used to control calcium and phosphate metabolism.

It is also known that such analogues are useful for inducing cell differentiation and for inhibiting undesired cell proliferation. For example, it is well recognized that vitamin D₃ produces 1α,25-dihydroxyvitamin D₃ (calcitriol) during normal metabolism. Calcitriol is a potent regulator of cell differentiation and proliferation as well as intestinal calcium and phosphorus absorption and bone calcium mobilization. Calcitriol is also known to affect

the immune system and this compound, as well as a variety of synthetic vitamin D₃ derivatives have been used in practical, clinical chemotherapy of such diverse human illnesses as osteoporosis, cancer, immunodeficiency syndromes and skin disorders such as dermatitis and psoriasis.

However, major research efforts are underway in an effort to prepare vitamin D₃ analogues as drugs in which calcitropic activity is effectively separated from cell growth regulation.

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Calcitriol may be structurally represented as follows:

The upper and lower ring portions of calcitriol may be called, for ease of reference, the C/D-ring and A-ring, respectively.

DESCRIPTION OF THE RELATED ART

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Especially for post-menopausal women, osteoporosis is a very serious illness that causes physical deformity and high susceptibility to bone (e.g., hip) fractures. In the general population aged above 65 years, osteoporosis ranks third to heart disease and cancer in terms of prevalence1. It is estimated that 30% of women at 75 years and 40% of women at 85 years have abnormal bone loss2. Calcitriol is being used, especially in Japan where dietary intake of calcium is low, for treatment of osteoporosis3. The Chugai Pharmaceutical Company has developed ED-71 (1) as a synthetic derivative of calcitriol, having a batter therapeutic index than calcitriol4. This 28-(3'-hydroxypropyloxy)-calcitriol has a two-fold stronger binding affinity to the rat plasma vitamin D-binding protein (DBP) than does calcitriol, suggesting that it circulates in the plasma with a longer half-life than calcitriol4. Furthermore, in animal models with osteoporosis, ED-71(1) is more effective than calcitriol4.

To probe structure-medicinal activity relationships in the hope of preparing a new osteoporosis drug with an even better therapeutic index than ED-71(1), we targeted the 2-carbanalogues 2 and 3, structurally represented below.

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These analogues were chosen because replacing one oxygen atom by a methylene group, a relatively small change in a large steroid molecule, was anticipated, based on the current working model for receptor binding of ED-71(1)4, not to interfere with such critical receptor binding. also, these analogues were chosen for chemical reasons, probing whether recently developed Diels-Alder methodology using 2-pyrones and monosubstituted alkenes could be extended to 1,2-disubstituted alkene dienophiles with reliable and faithful transfer of olefin geometry ultimately into the stereochemical relationships at the 1- and 2positions of the steroid targets. We record here the results of these chemical explorations and biological evaluations.

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SUMMARY OF THE INVENTION

The present invention is for a vitamin D_3 analogue which includes a 2-substituted alcohol or fluoride. The preferred alcohol substituent is exemplified by the structural formula -(CH₂)₄OH and the preferred fluoride substituents by the structural formulas -(CH₂)₄F.

The preferred diastereomers of the 2-substituted -(CH₂)₄OH are represented by the following structural formulas:

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The present invention is also for the related method of preparation of a vitamin D_3 analogue which includes a 2-substituted alcohol or fluoride starting with 2+4-cycloaddition of commercially available methyl 2-pyrone-3-carboxylate.

- 5 -

Diastereomeric 2-substituted calcitriol analogues were prepared in only eleven chemical operations, starting with 2+4-cycloaddition of commercially available methyl 2-pyrone-3-carboxylate. Highlights of this 5 convergent and stereo controlled synthetic approach are as follows: (1) retention of reactant dienophile geometry in the product bicyclic lactone, characteristic of a concerted inverse-electron-demand Diels-Alder cycloaddition; 10 (2) an improved decarboxylation procedure involving chemospecific allyloxide opening of a lactone ring in the presence of a methyl ester and then non-high pressure palladium-promoted allylic ester decarboxylation; and (3) use of the 15 enantiomerically pure C,D-ring chiron 14 to resolve racemic A-ring phosphine oxide 13.

DETAILED DESCRIPTION OF THE PRESENTLY PREFERRED EXEMPLARY EMBODIMENTS

20 Uncatalyzed Diels-Alder cycloadditions
between highly polarized dienes and dienophiles
can occur step-wise rather than in the usual
concerted fashion⁶. To probe this issue within the
context of inverse-electron-demand Diels-Alder
25 cycloadditions using 2-pyrones substituted at the
3-position with highly electron-withdrawing (e.g.

3-sulfonyl, 3-acyl) substituents⁵, 3-ptoluenesulfonyl-2-pyrone and 1,2-disubstituted
alkenes 4E and separately 4Z were placed under
high pressure. In both of these electronically
matched cases, the electron-poor 2-pyrone diene
and the electron-rich vinylic ether cycloadded to
produce isolable bicyclic lactone adducts without
undesirable and often-encountered extrusion of CO₂
(Scheme I)

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The most important aspect of these Diels-Alder reactions from a mechanistic viewpoint is that the cycloadducts retained the stereochemical information in the reactant vinylic ethers: dienophile 4E led exclusively to trans-4,5oriented products 5a and 5b, whereas dienophile 4Z led exclusively to cis-4,5-oriented products 6a Therefore, these polarized 2+4cycloadditions must occur in a concerted rather than in a step-wise fashion6. The assignments of the 4,5-positional relationships were based on extensive precedent⁵, and the assignments of the 4,5-stereochemical relationships were based on the match of the 400 MHz ¹H NMR J_{4,5} coupling constants with those calculated using the Karplus equation for energy-minimized structures generated using Chem-3D (Scheme I) 7. Bicyclic lactone 6a, the very major cycloadduct, differed in a characteristic way⁵ from bicyclic lactone 6b in terms of the

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chemical shift of the bridgehead hydrogen atom (δ 4.98 vs. 5.04) and the chemical shift of the C_{λ} hydrogen atom (δ 4.55 vs. 3.83). Also, irradiation of the C_5 hydrogen atom of lactone 6acaused an 11 % noe on the C4 hydrogen atom. large discrepancy between the observed and the calculated $J_{4.5}$ coupling constant for cis-4.5disubstituted bicyclic adduct 6b was of considerable concern. Therefore, a series of similar cis-4.5-disubstituted bicyclic lactones was prepared. Examination of their $J_{4.5}$ coupling constants (Table I) showed a very subtle effect of the nature of the substituents on this magnitude of the vicinal coupling constant. For example, the cycloadduct derived from the 6-membered cyclic vinyl ether showed $J_{4.5}=0$ Hz (entry 2), whereas that derived from the 5-membered cyclic vinyl ether showed $J_{4.5}=8.5$ Hz (entry 3). Finally, based on literature analogies in which E-alkenes underwent 2+4-cycloadditions more easily than Zalkenes6e-9, we were surprised to find that vinylic ether geometric isomer 42 reacted considerably faster than isomer 4E with 3-tolysufonyl-2-pyrone. Whereas vinylic ether isomer 42 yielded almost exclusively the endo-cycloadduct 6a, as expected from previous results⁵, vinylic ether isomer 4E gave a 1:2 mixture of cycloadducts 5a:5b.

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Prom a synthesis viewpoint, cycloadducts 5 and 6 turned out to be disappointing. Despite close structural similarity with other bridgeheadsubstituted toluenesulfonyl cycloadducts and related cyclohexene systems that underwent smooth reductive-desulfonylation8, we were unsuccessful using a variety of conditions (e.g., Al-Hg, Na-Hg, Raney nickel, Li/NH3) to effect high-yielding reductive-desulfonylation. Also, it was anticipated that ultimate conversion of the methyl ether functionality into the desired alcohol group would be more difficult than deprotection of a silyl ether. Therefore, silylated vinylic ether 72, prepared according to literature precedent as illustrated in Scheme II9, and commercially available methyl 2-pyrone-3-carboxylate were subjected to high pressure cycloaddition (eq. 1).

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Bicycloadduct 8 was the major product, isolated on gram scale in 60% yield, with the oxygen substituted at position-4, as expected based on the polar nature of the Diels-Alder cycloaddition and also on literature precedent⁵, with a cis-4,5-stereochemical relationship. This stereochemical outcome was expected based on the results in Scheme I with the 3-sulfonyl-2-pyrone and was confirmed by the observed large ¹H NMR J_{4.5} coupling constant (8.6 Hz)⁷ and by the characteristic chemical shifts of the bridgehead

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hydrogen atom at δ 5.1 and the C_4 hydrogen atom at δ 4.7. Also, irradiation of the C_5 hydrogen atom caused a 14% nOe on the C_4 hydrogen atom, confirming their cis-4,5-relationship. We have not succeeded in preparing cleanly the isomeric silylated vinylic ether 7E for cycloaddition with methyl 2-pyrone-3-carboxylate.

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Having served its chemical function of activating the pyrone diene (the unsubstituted parent 2-pyrone is unreactive) 5d,10 for cycloaddition with the electron-rich vinylic ether 72, the bridgehead carboxylate ester group in bicyclic lactone adduct 8 had to be removed. complement our two-step lactone methanolysis and high-pressure procedure for this type of decarboxylation11, we report now a new and more convenience (i.e., not high pressure) two-step protocol (Scheme III). Although bicyclic lactone methyl ester 8 has two ester carbonyl groups, it was gratifying to find that the lactone ring was chemospecifically attacked by lithium allyloxide to produce mixed methyl allyl malonate 9 in 75% yield. In accord with literature precedent 12, allyl ester 9 was smoothly decarboxylated using palladium acetate; an unexpected but desired benefit of this procedure was conjugation of the cyclohexene double bond, giving the contiguously

SCHEME II.

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tetrasubstituted cyclohexene 10 in 92% yield (Scheme III).

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Without any surprises, highly functionalized cyclohexene alcohol ester 10 was O-silylated and then reduced to form allylic alcohol 11 (Scheme IV). One flask Claisen rearrangement followed by spontaneous thermal sulfoxide elimination was achieved using our sulfinylated ortho ester protocol giving an unusually favorable >10:1 Z:E ratio of dienoate esters from which the desired Z-dienoate 12 was isolated by chromatography in 80% yield.

Established reactions as outlined in Scheme

IV below provided the crucial, fully O-protected,
racemic, A-ring phosphine oxide 13. Lythgoe-type
coupling 14 of the conjugate base of phosphine oxide
13, generated using phenyllithium 15, with C,D-ring
ketone 14 of natural absolute configuration
produced O-silylated derivatives of diastereomeric
2-(4'-hydroxylated) calcitriol analogues 3 and 3',
isolated in 50% yield.

Fluoride-induced cleavage of the silyl protecting groups proceeded easily at three of the four silylated alcohol groups; desilylation at the C₁ secondary alcohol position, however, was unexpectedly slow, requiring considerably more vigorous reaction conditions.

In another context 16 , we have observed that the C_1 secondary alcohol unit in calcitriol is chemically less reactive than the C_3 secondary alcohol toward esterifying reagents.

Nevertheless, fluoride-assisted quadruple desilylation under more vigorous conditions eventually yielded the desired calcitriol analogues 3 and 3.

in enantiomerically pure form. Tentative
assignments of stereochemistry to diastereomers 3
and 3' were made by H NMR in analogy with closely
related calcitriol analogues; in diastereomeric
pairs differing only by inversions of
stereochemistry at positions 1-3 but not in the
C,D-ring or in the side chain, the 1α-substituted
diastereomer characteristically showed a lower
field absorption for the C₁₈-methyl group and for
one proton of the C₁₉-methylene group (Table II).

						7			l
3′	ω								
18-0H	1α-OH	в-сн,сн,он	α-СН ₂ СН ₂ ОН	в-сн ₂ он	α-CH ₂ OH	в-он	·α-0H	1	
2B-(CH ₂₎ OH	2α-(CH ₂),OH	ı	1	ţ		в-он	α-0H	2	
α-ОН	в-он	.α-0H	в-он	α-ОН	в-он	α-ОН	в-он	ω.	
0.52	0.54	0.51	0.54	0.50	0.54	0.53	0.54	CIB	
4.98	5.00	4.88	4.90	4.99	5.02	5.11	5.14	C ₁₀	
work	this	13b	13b	15	15	17	17	Ref	

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Thus, in only 11 steps from commercially available methyl 2-pyrone-3-carboxylate, two new vitamin D_3 analogues have been prepared, and the C,D-ring chiron 14 has been used to resolve racemic A-ring phosphine oxide 13.

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Preliminary biological evaluation of synthetic diastereomers 3 and 3' involved measuring their relative binding affinities to rat vitamin D binding protein (DBP) and also to the vitamin D receptor of bovine thymus (Table III). 5 Diastereomer 3 had significantly higher affinity than ED-71 (1) for the vitamin D binding protein, but it had extremely low affinity for the vitamin D receptor; whether this separation of binding affinities has important mechanistic and/or 10 medicinal value remains to be established. Diastereomer 3', on the other hand, had lower affinity than ED-71 (1) for the vitamin D binding protein, but it had higher affinity than ED-71 (1) for the vitamin D receptor. Determining the 15 impact of these differences on possible use of these new analogues for chemotherapy of osteoporosis requires further biological testing.

TABLE III.

20 RELATIVE BINDING AFFINITIES

		DBP	D RECEPTOR
	25 (OH) -D ₃	90	
	1,25(OH) ₂ -D ₃	1	1 .
	Ed-71(1)	3.7	0.21
25	· 3	5.0	0.01
	31	1.9	0.28

Relative binding affinities of diastereomers

3 and 3' to the vitamin D binding protein and to
the vitamin D receptor showed some unusual trends.

Diastereomer 3' surprisingly bound much more
effectively to the vitamin D receptor than did the
established osteoporosis drug candidate ED-71 (1).

EXAMPLES

General

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Tetrahydrofuran and diethyl ether were distilled from benzophenone ketyl prior to use. 10 Methylene chloride and triethylamine were distilled from calcium hydride immediately prior to use. Commercially available anhydrous solvents were used in other instances. All reagents were purchased from Aldrich Chemical Co. (Milwaukee, 15 WI) and unless otherwise specified were used as received without further purification. All reactions were run in flame-dried flasks under nitrogen unless otherwise specified. FT-IR spectra were determined using a Perkin-Elmer Model 20 1600 FT-IR spectrophotometer. The ¹H NMR were recorded on a Varian XL-400 spectrometer and Bruker AMX-300 spectrometer operating at 400 and 300 MHz, respectively. The 13C spectra were recorded on the same instruments operating at 100 25 and 75 MHz, respectively. High resolution mass spectra were obtained on a two sector-high-

resolution VG-70S mass spectrometer run at 70 eV.

A Leco Corp. Model No. PG-200-HPC 13 Kbar

apparatus was used for the high-pressure studies.

Silylated Vinylic Ether 72

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To a 100 mL round-bottomed flask containing 2.1 g (17.5 mmol) of 1,6-hexanediol, 2.7mL (19.2 mmol) of triethylamine, and 10 mg of N,N-dimethylaminopyridine in 35 mL CH₂Cl₂ was added 5 mL (19.2 mmol) of t-butylchlorodiphenylsilane. The reaction was stirred at room temperature for 12 h, or until complete by TLC. The reaction was quenched with 10 mL water, the organic layer was separated, and the aqueous layer was washed with CH₂Cl₂. The combined organic layers were dried over MgSO₄, filtered, and concentrated. The crude product was purified by column chromatography (10-20% EtOAc/Hexane) to afford 3.73 g (10.45 mmol, 60% yield) of the monosilated product as a clear oil.

To a 250 mL round-bottomed flask containing 10.14 mL (125.4 mmol) of pyridine in 50 mL of CH₂Cl₂ was slowly added 6.27 g (62.7 mmol) of chromium trioxide. The resultant deep burgundy solution was stirred for 15 minutes at room temperature. At the end of this period, a solution of the above monosilated diol in 50 mL CH₂Cl₂ was added via cannula. A tarry, black

deposit separated immediately. This mixture was stirred for 1h at which time the CH₂Cl₂ was removed on a rotary evaporator and the tar was diluted with Et₂O. This heterogeneous mixture was filtered through silica gel to give a yellow liquid which was concentrated. The crude product was purified by column chromatography (5% EtOAc/Hexane) to give 3.17 g (8.9 mmol, 85% yield) of the aldehyde as a clear oil. ¹H NMR (300 MHz, CDCl₃) & 9.74 (s, 1H), 7.675 (m, 4H), 7.40 (m, 6H), 3.66 (t, J=6.2 Hz, 2H), 2.34 (m, 2H), 1.58 (m, 4H), 1.41 (m, 2H), 1.04 (s, 9H), IR (CHCl₃) 3013 cm⁻¹, 2931 cm⁻¹, 1713 cm⁻¹.

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The above aldehyde was then diluted with 20 mL of benzene and 1.64 mL (11.75 mmol) of 15 triethylamine. The resulting mixture was cooled to 0°C and 2.45 ml (10.68 mmol) of tbutyldimethylsilyl triflate was added. The reaction was allowed to warm to room temperature and stirred for 1 h. The reaction was quenched 20 with brine and diluted with Et,O. The organic layer was separated, and the aqueous layer was washed with Et20. The combined organic layers were dried over MgSO4, filtered, and concentrated. crude product was purified by column 25 chromatography (Hexane) or silica gel that was slurry-packed with 1% triethylamine/hexane. silylated vinylic ether 72 (1.67 g, 3.56 mmol, 40%

yield) was a clear oil; $R_f=0.8$ (25% EtOAc/Hexane); 1 H NMR (300 MHz, CDCl₃) δ 7.65 (m, 4H), 7.41 (m, 6H), 6.17 (dt, J=5.9, 1.5 Hz, 1H), 4.43 (dd, J=7.4, 5.9 Hz, 1H), 3.61 (t, J=6.6 Hz, 2H), 2.09 (ddt, J=14.7, 7.4, 1.5 Hz, 2H), 1.54 (m, 2H), 1.37 (m, 2H), 0.92 (s, 9H), 0.12 (s, 6H); 13 C NMR (75 MHz, CDCl₃) δ 138.50, 135.54 (4), 134.15 (2), 129.50, 129.43, 127.55 (2), 127.53, 110.54, 63.93, 32.31, 26.87 (3), 25.99 (3), 25.80, 25.66, 23.36, 19.22, -2.92, -5.36; IR (CHCl₃) 3013, 2931, 1654, 1108 cm⁻¹; HRMS m/e calcd. for $C_{24}H_{35}O_{2}Si_{2}$ 411.2176, found 411.2179.

Cycloadduct 8

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A 12 cm piece of 3/8" heat shrinkable teflon tubing (Ace Glass cat. #12685-40) was sealed on 15 one end with a glass dowel plug by using a heat gun. To this 500.0 mg (3.24 mmol) of methyl 2pyrone-3-carboxylate (Aldrich), 2g (4.26 mmol) of silylated vinylic ether 72, 10 mg of barium carbonate, and 2 mL of dry CH2Cl2 was added. 20 open end of tubing was then sealed in a similar fashion with a second glass dowel plug. This 'sealed tube' was the pressurized at 10-11 Kbar at room temperature for 4 days. The reaction mixture was concentrated on a rotary evaporator and the 25 residue was purified by column chromatography (5% EtOAc/Hexane) to give 1.21 g (1.94 mmol, 60%) of

the cycloadduct 8 as a clear oil; R_f=0.58 (25% EtOAc/Hexane); 1 H NMR (300 MHz, CDCl₃) δ 7.65 (m, 4H), 7.41 (m, 6H), 6.78 (dd, J=7.8, 2.8 Hz, 1H), 6.46 (dd, J=7.8, 5.1 Hz, 1H), 5.09 (ddd, J=5.1, 3.8, 1.0 Hz, 1H), 4.71 (dd, J=7.6, 1.00 Hz, 1H), 5 3.89 (s, 1H), 3.66 (t, J=6.1 Hz, 3H), 2.35 (ddd, J=7.6, 3.8, 2.8 Hz, 1H), 1.56 (m, 2H), 1.26 (m, 2H), 1.06 (s, 9H), 0.91 (m, 2H), 0.78 (s, 9H), 0.012 (s, 3H), 0.006 (s, 3H); ¹³C NMR (75 MHz, CDCl₃) & 168.93, 167.51, 135.51 (4), 133.70 (2), 10 130.46, 129.53 (2), 128.64, 127.54 (4), 76.12, 69.36, 63.12, 52.7, 44.72, 32.37, 27.09, 26.75 (3), 25.64 (3), 25.56, 25.31, 23.34, 19.09, 18.04, -3.90, -5.03, IR (CHCl_x) 1760, 1743 cm⁻¹; HRMS m/e calcd. for $C_{31}H_{41}O_6Si_2$ 565.2442, found 565.2450. 15

Mixed Malonate 9

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To a 25 mL round-bottomed flask with 500 mg (0.80 mmol) of cycloadduct 8 and 2 mL of CH₂Cl₂ at 0°C was added dropwise, via syringe, 962 µL of a freshly made 1.0 M solution of lithium allyloxide in allyl alcohol. The reaction mixture was allowed to warm to room temperature after the addition. Reaction was complete by TLC after 2 hours. The mixture was quenched with 2 mL aq. NH₄Cl and extracted with CH₂Cl₂. The organic layer was dried with MgSO₄, filtered, and concentrated on a rotary evaporator. The residue was purified by

column chromatography (0-20% EtOAc/Hexane) to give 410 mg (0.60 mmol, 75%) of the desired malonate as a clear oil; R=0.42 (25% EtOAc/Hexane); H NMR (300 MHz, CDCl₃) δ 7.68 (m, 4H), 7.41 (m, 6H), 5.96 (s, 2H), 5.83 (ddt. J=17.2, 10.5, 5.6 Hz, 1H), 5 5.36 (dd, J=17.2, 1.5 Hz, 1H), 5.22 (dd, J=10.5, 1.5 Hz, 1H), 4.76 (s, 1H), 4.63 (ddt, J=13.3, 5.6, 1.4 Hz, 1H), 4.51 (ddt, J=13.3, 5.6, 1.4 Hz, 1H), 4.02 (t, J=8.6 Hz, 1H), 3.74 (s, 3H), 3.69 (t, J=6.1 Hz, 2H), 1.71 (m, 2H), 1.64 (m, 2H), 1.57 10 (s, 1H), 1.28 (m, 2H), 1.05 (s, 9H), 0.81 (s, 9H), 0.07 (s, 3H), -0.06 (s, 3H); 13 C NMR (75 MHz, $CDCl_3$) δ 169.11, 167.95, 135.41 (4), 134.07 (2), 133.89, 131.17, 129.34 (2), 127.44 (4), 122.47, 118.43, 71.70, 68.64, 65.72, 63.89, 61.38, 52.57, 15 46.04, 32.45, 26.76 (3), 25.94 (3), 23.25, 19.04, 18.27, 14.02, -3.79, -4.44; IR (CHCl₃) 1737 cm⁻¹; HRMS m/e calcd. for $C_{34}H_{47}O_{7}Si_{2}$ 623.2860, found 623.2865.

To a 25 mL round-bottomed flask with 380 mg (0.56 mmol) of the allyl ester alcohol, 78 μL (0.67 mmol) of 2,6-lutidine, and 1.5 mL of CH₂Cl₂ at 0°C was added 154 μL (0.67 mmol) of t-butyldimethylsilyl trifluoroethanesulfonate dropwise via syringe. The reaction was complete by TLC after 10 minutes. The reaction was quenched at 0°C with 1 mL water and allowed to warm to room temperature. Extraction with CH₂Cl₂

followed by drying with MgSO, filtration, and concentration afforded a viscous oil which was purified by column chromatography (5% EtOAc/Hexane) giving 390 mg (0.49 mmol, 87%) of 0silylated mixed malonate 9 as a clear oil; R,=0.6 5 (25% EtOAc/Hexane); ¹H NMR (300 MHz, CDCl₃) δ 7.68 (m, 4H), 7.41 (m, 6H), 5.87 (d, J=2.68 Hz, 2H), 5.83 (ddt, J=17.2, 10.5, 5.6 Hz, 1H), 5.36 (dd, J=17.2, 1.5 Hz, 1H), 5.22 (dd, J=10.5, 1.5 Hz, 1H), 4.76 (s, 1H), 4.63 (ddt, J=13.3, 5.6, 1.4 Hz, 10 1H), 4.51 (ddt, J=13.3, 5.6, 1.4 Hz, 1H), 4.13 (d, J=9.2 Hz, 1H), 3.74 (s, 3H), 3.69 (t, J=6.1 Hz, 2H), 1.76 (m, 2H), 1.61 (m, 2H), 1.55 (s, 1H), 1.28 (m, 2H), 1.06 (s, 9H), 0.88 (s, 9H), 0.83 (s, 9H), 0.08 (s, 3H), 0.06 (s, 3H), 0.04 (s, 3H), -15 0.06 (s, 3H); 13 C NMR (75 MHz, CDCl₃) δ 169.27, 167.96, 135.45 (4), 133.99 (2), 133.97, 131.33, 129.38 (2), 127.48 (4), 121.83, 118,43, 71.70, 69.16, 65.69, 64.00, 61.47, 52.52, 46.10, 32.67, 26.81 (3), 25.99 (3), 25.80 (3), 23.25, 19.10, 20 18.35, 17.99, 14.11, -3.83, -4.20, -4.34, -4.70; IR (CHCl₃) 1737 cm⁻¹; HRMS m/e calcd. for C₄₀H₆₁O₇Si₃ 737.3725, found 737.33730.

Cyclohexene Ester 10

25 A mixture of 380 mg (0.48 mmol) of malonate 9, 23 μ L (0.60 mmol) formic acid, 87 μ L (0.62 mmol) triethylamine, 10 mg (0.04 mmol)

triphenylphosphine, and 2 mg (0.01 mmol) palladium acetate in 1.5 mL dioxane was sealed in a 5 mL hydrolysis tube and heated at 100°C for 12 h. After evaporation of dioxane, 1 N HCl (lmL) was added, and the mixture was extracted with CH2Cl2 (2 5 $mL \times 2$). The organic solution was washed with saturated NaHCO3 and dried over MgSO4, filtered, and concentrated. The oily residue was purified by column chromatography (0-10% EtOAc/Hexane) to give 310 mg (0.44 mmol, 92%) of cyclohexene ester 10 10 as a clear oil: R_f=0.50 (25% EtOAc/Hexane); ¹H NMR (300 MHz, CDCl₃) δ 7.68 (m, 4H), 7.41 (m, 6H), 6.87 (t, J=1.5 Hz, 1H), 5.76 (s, 1H), 4.72 (s, 1H), 4.16 (m, 1H), 3.99 (m, 1H), 3.74 (s, 3H), 3.69 (t, J=6.1 Hz, 2H), 2.60 (dt, J=14.3, 5.6 Hz, 15 1H), 2.10 (ddd, J=16.9, 8.4, 1.5 Hz, 1H), 1.58 (m, 2H), 1.55 (s, 1H), 1.38 (m, 2H), 1.26 (m, 2H), 1.05 (s, 9H), 0.89 (s, 9H), 0.83 (s, 9H), 0.05 (s, 3H), 0.04 (s, 3H), -0.07 (s, 3H), -0.09 (s, 3H); 13 C NMR (75 MHz, CDCl₃) δ 167.01, 135.51 (4), 20 134.05 (2), 133.16, 132.35, 129.45 (2), 127.55 (4), 66.89, 64.16, 51.50, 47.43, 33.1, 26.87, 25.95, 25.92, 25.89, 25.84, 25.68, 23.31, 19.17, 18.42, 18.06, -4.09, -4.37, -4.74, -5.39; IR (CHCl₃) 1713 cm⁻¹; HRMS m/e calcd. for $C_{36}H_{57}O_5Si_3$ 25 653.3514, found 653.3516.

Allylic Alcohol 11

To a 25 mL round-bottomed flask containing 695 mg (0.98 mmol) of cyclohexene ester 10 in 8 mL toluene at -78°C was added 2.15 ml (2.15 mmol) of 1 M diisobutylaluminum hydride in toluene dropwise 5 via syringe. The reaction was complete by TLC after 1h. The reaction was quenched by addition at -78°C of 5 mL of 2 M potassium sodium tartrate followed by dilution with 10 mL EtOAc. After the mixture was allowed to warm to room temperature 10 and stirred for 0.5 h, two layers were visible. These were separated and the aqueous layer was washed with EtOAc. The combined organic fractions were washed with water, and brine, and dried over MgSO,, filtered, and concentrated. The residue was 15 purified by column chromatography (0-25% EtOAc/Hexane) to give 518 mg (0.76 mmol, 77%) of allylic alcohol 11 as a clear oil: R,=0.35 (25% EtOAc/Hexane); 1 H NMR (300 MHz, CDCl₃) δ 7.68 (m, 4H), 7.41 (m, 6H), 5.51 (t, J=1.5 Hz, 1H), 4.80 20 (s, 1H), 4.22 (d, J=12.0 Hz, 1H), 4.16 (m, 1H), 3.99 (m, 1H), 3.74 (s, 3H), 3.69 (t, J=6.1 Hz,2H), 2.60 (dt, J=14.3, 5.6 Hz, 1H), 2.10 (ddd, J=16.9, 8.4, 1.5 Hz, 1H), 1.58 (m, 2H), 1.55 (s, 1H), 1.38 (m, 2H), 1.26 (m, 2H), 1.05 (s, 9H), 25 0.89 (s, 9H), 0.83 (s, 9H), 0.05 (s, 3H), 0.04 (s, 3H), -0.07 (s, 3H), -0.09 (s, 3H); 13 C NMR (100 MHz, CDCl₃) δ 137.81, 135.51 (4), 134.04 (2),

129.47 (2), 127.55 (4), 122.65, 70.03, 68.32, 65.68, 63.77, 46.36, 33.11, 31.28, 26.85 (3), 25.90 (3), 25.82 (3), 25.51, 25.43, 24.52, 24.03, 19.21, 18.06, -4.20, -4.82 (2), -4.92; IR (CHCl₃) 3401, 1713 cm⁻¹; HRMS m/e calcd. for $C_{34}H_{57}O_{4}Si_{3}$ 625.3565, found 625.3570.

z-Dienoate 12

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To 510 mg (0.75 mmol) of allylic alcohol 11 in a sealable hydrolysis tube was added 642 mg (2.2 mmol) triethylphenyl sulfinyl orthoacetate 10 and 1.0 mg 2,4,6-trimethylbenzoic acid and 2 mL CH2Cl2. The tube was purged with argon and sealed, then heated at 100°C for 12h. The reaction mixture was concentrated. Crude ¹H NMR showed a >10:1 mixture of Z:E isomers. The crude product 15 was purified via PTLC (10%EtOAc/Hexane) to give 450 mg (0.60 mmol, 80%) of Z dienoate 12 as a clear oil; R=0.61 (25% EtOAc/Hexane); 1H NMR (400 MHz, CDCl₃) δ 7.68 (m, 4H), 7.41 (m, 6H), 5.61 (s, 1H), 5.17 (t, J=1.8 Hz, 1H), 5.09 (t, J=1.3 Hz, 20 1H) 4.55 (dd, J=5.5, 1.3 Hz, 2H), 4.12 (q, J=7.2 Hz, 2H), 3.97 (t, J=3.8 Hz, 1H), 3.66 (t, J=6.1Hz, 2H), 2.48 (dd, J=5.5, 1.4 Hz, 1H), 2.15 (dd, J=3.8, 1.7 Hz, 1H), 1.70 (m, 1H), 1.58 (m, 2H), 1.55 (s, 1H), 1.38 (m, 2H), 1.26 (m, 2H), 1.04 (s, 25 9H), 0.87 (s, 18H), 0.05 (s, 6H), 0.04 (s, 6H); ¹³C NMR (75 MHz, CDCl₃) δ 165.92, 157.09, 153.29,

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145.39, 135.52 (4), 134.09 (2), 129.47 (2), 127.55 (4), 117.17, 70.23, 63.91, 59.67, 50.54, 32.98, 26.86 (3), 25.80 (3), 25.74 (3), 19.20, 18.18, 17.98, 14.30, 14.13, -4.66, -4.83, -4.87, -5.17; IR (CHCl₃) 3025, 2931, 1713 cm⁻¹; HRMS m/e calcd. for $C_{43}H_{70}O_5Si_3$ 750.4531, found 750.4538.

Phosphine Oxide 13

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To a 25 mL round-bottomed flask containing 180 mg (0.24 mmol) of dienoate 12 in 2 mL toluene at -78°C was added dropwise 530 μ L (0.53 mmol) of a 1 M solution of diisobutylaluminum hydride via syringe. After 1h the reaction was complete by TLC. The reaction was quenched at -78°C by addition of 2 mL of 2 N potassium sodium tartrate and dilution with 5 mL EtOAc. The mixture was allowed to warm to room temperature and stirred 0.5 h until two distinct phases appeared. organic phase was separated, and the aqueous phase was washed with EtOAc. The combined organic extracts were washed with water and brine, dried over MgSO4, filtered, and concentrated. The crude mixture was quickly purified by column chromatography (10-20% EtOAc/Hexane) to give 167 mg (0.24 mmol) of the desired allylic alcohol.

To a 10 mL round-bottomed flask containing 152 mg (1.14 mmol) N-chlorosuccinimide in 3 mL CH₂Cl₂ at 0°C was added 90 mL (1.22 mmol) of

dimethyl sulfide via syringe. A white precipitate formed immediately upon addition. This mixture was cooled to -20°C and stirred for 20 minutes. The above allylic alcohol in 2 mL CH₂Cl₂ was then added to the heterogeneous mixture via cannula. The reaction was stirred for 0.5 h at -20°C and then allowed to warm to room temperature and stirred for an additional 1 h. The organic layer was washed with water, brine, dried with MgSO₄, filtered, and concentrated. The crude product was passed through florisil (5% EtOAc/Hexane) to afford 154 mg (0.21 mmol) of the desired allylic chloride as a yellow oil.

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To a 10 mL round-bottomed flask containing the above allylic chloride in 2 mL THF at -78°C was added dropwise via cannula a 0.5 M solution of potassium diphenylphosphide in THF. The addition was stopped once the red color persisted. After 1 h the reaction was allowed to slowly warm to 0°C at which point it was complete by TLC. The reaction was quenched with 2 drops of water and the THF was removed. The residue was diluted with 2 mL CH₂Cl₂ and 10 drops of 30% H₂O₂ was added. After 1 h the reaction was diluted with CH₂Cl₂ and water and the layers were separated. The organic phase was concentrated. The crude product was purified by column chromatography (25-50%) EtOAc/Hexane) to give 121 mg (0.13 mmol, 54% from

Z-dienoate 12) of the phosphine oxide 13 as a clear oil: $R_f=0.56$ (75% EtOAc/Hexane); ¹H NMR (400 MHz, $CDCl_3$) & 7.74-7.36 (m, 20H), 5.3 (dd, J=15.1Hz, 6.6H), 5.11 (s, 1H), 4.75 (s, 1H), 4.42 (d, J=3.2 Hz, 1H), 3.87 (dd, J=8.9, 4.9 Hz, 1H), 3.65 5 (t, J=6.6 Hz, 3H), 3.40 (dt, J=15.1, 8.9 Hz, 1H), 3.14 (dt, J=16.0, 6.7 Hz, 1H), 2.4 (dd, J=13.3, 3.2 Hz, 1H), 2.06 (dd, J=14.2, 2.9 Hz, 1H), 1.56 (m, 1H), 1.53 (m, 2H), 1.43 (m, 2H), 1.35 (m, 2H), 1.04 (s, 9H), 0.88 (s, 9H), 0.82 (s, 9H), 0.02 (s, 10 3H), 0.01 (s, 3H), -0.01 (s, 3H), -0.03 (s, 3H); 13 C NMR (75 MHz, CDCl₃) δ 146.20 (2), 141.17, 135.50 (4), 134.06 (2), 131.71, 131.04, 130.92 (4), 129.45 (2), 128.63 (4), 128.47 (2), 127.53 (4), 114.39, 70.15, 63.90, 50.72, 32.97, 30.09, 15 26.83 (3), 25.79 (3), 25.05, 24.06, 19.18, 18.04, -4.58, -4.69, -4.89, -5.07; IR (CHCl₃) 3025, 2954, 1472, 1255 cm⁻¹; HRMS m/e calcd. for C40H48O4Si3 835.4163, found 835.4169.

20 Calcitriol Analogues 3 and 3'

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To a 10 mL round-bottomed flask containing 95 mg (0.103 mmol) of phosphine oxide 13 in 1.4 mL of THF at -78°C was added dropwise 67 μ L (0.103 mmol) of a 1.54 M solution of phenyllithium in THF. The resultant red solution was allowed to stir for 10 minutes. A solution of 20 mg (0.056 mmol) of C-D ring 14 in 1 mL of THF was added via cannula. The

reaction was complete after 1 h by TLC following disappearance of C-D ring. The reaction was quenched by addition of 1 mL 1:1 KHCO3/2 N potassium sodium tartrate. The layers were separated and the organic phase was washed with 5 brine, dried over MgSO, filtered and concentrated. The crude product was rapidly purified by filtration through silica gel using 50% EtOAc/Hexane as solvent to give 47 mg (0.052 mmol, 50% yield) of the coupled product. The silyl 10 ethers were cleaved by redissolving the product in 1 mL of THF and treating with 220 μ L (0.22 mmol) of tetrabutylammonium fluoride. After 24 h the reaction was diluted with water and the layers were separated, dried over MgSO, filtered and 15 concentrated. The diastereomers were separated and purified by reverse phase HPLC (30-20% H,O/CH,CN on a C-18 semi-prep. column) to afford 1.1 mg (0.002 mmol, 4% yield) of analogue 3, and 4.4 mg (0.008 mmol, 16% yield) of analogue 31, 20 both as white solids. 1H NMR of 3 (300 MHz, CDCl₃) δ 6.41 (d, J=11.6 Hz, 1H), 6.02 (d, J=11.6 Hz, 1H), 5.29 (s, 1H), 5.02 (s, 1H), 4.35 (s, 1H), 3.82 (m, 1H), 3.68 (t, J=6.1 Hz, 2H), 2.78 (d, J=11.5 Hz, 1H), 2.62 (dd, J=11.5, 3.5 Hz, 1H), 25 2.20 (dd, J=10.2, 3.5 Hz, 1H), 1.98 (m, 2H), 1.68-1.20 (m), 0.92 (s, 3H), 0.90 (s, 3H), 0.54 (s, 3H). H NMR of 3' (300 MHz, CDCl₃) δ 6.40 (d,

J=11.6 Hz, 1H), 5.98 (d, J=11.6 Hz, 1H), 5.27 (s, 1H), 4.98 (s, 1H), 4.38 (s, 1H), 3.87 (m, 1H), 3.68 (t, J=6.1 Hz, 2H), 2.82 (d, J=11.5 Hz, 1H), 2.65 (dd, J=11.5, 3.5 Hz, 1H), 2.24 (dd, J=10.2, 3.5 Hz, 1H), 1.68-1.20 (m), 0.92 (s, 3H), 0.90 (s, 3H), 0.52 (s, 3H); UV-Vis (MeOH) λ_{max} 268 nm (ϵ 15,600); HRMS m/e calcd. for $C_{31}H_{50}O_{3}$ 470.3760, found 470.3771.

2-Fluorides

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The above-described methods for producing the vitamin D₃ analogues with alcohols substituted in the 2-position can be readily modified to produce comparable analogues with fluorides substituted in the two position (18-21). Table IV lists the fluoride analogues of the present invention.

TABLE IV

2-SUBSTITUTED FLUORIDES OF CALCITRIOL

Position

<u>3</u> .	<u>2</u>	<u>1</u>
3 <i>β</i>	2α -(CH ₂) ₄ F	1α
3α	2β -(CH ₂) ₄ F	1 β .
3 <i>β</i>	2α -(CH ₂) ₃ F	10
3α	2β -(CH ₂) ₃ F	1β

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- 20 While the invention has been described in connection with what is presently considered to be the most practical and preferred embodiments, it is to be understood that the invention is not to be limited to the disclosed embodiments, but on the contrary is intended to cover various modifications and equivalent arrangements included within the spirit and scope of the appended claims.

WHAT IS CLAIMED IS:

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1. A vitamin D_3 analogue which includes a substituent in the 2-position selected from the group consisting of alcohols and fluorides.

- 2. The analogue of claim 1 wherein said substituent is $-(CH_2)_4OH$.
- 3. The analogue of claim 2 having the following structural formula:

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4. The analogue of claim 2 having the following structural formula:

- 5. The analogue of claim 1 wherein said substituent is $-(CH_2)_3F$.
- 6. The analogue of claim 1 wherein said substituent is $-(CH_2)_4F$.
 - 7. A method for the preparation of a vitamin D_3 analogue which includes an alcohol substituent in the 2-position, comprising the steps of:
 - a) subjecting methyl 2-pyrone-3-carboxylate and a silylated vinylic ether to high pressure cycloaddition to form a bicycloadduct;
 - b) reacting the bicycloadduct with lithium allyloxide to produce a mixed methyl allyl malonate;

c) decarboxylating the mixed methyl allyl malonate with palladium acetate to yield a cyclohexene ester;

- d) silylating and reducing the cyclohexene
 ester to form an alcohol;
- e) subjecting the allylic alcohol to a Claisen rearrangement followed by spontaneous thermal sulfoxide elimination to form a Z-dienoate;

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- f) reacting the Z-dienoate with a hydride to form an allylic alcohol;
 - g) reacting the allylic alcohol with Nchlorosuccinimide and dimethyl sulfide to form an allylic chloride;
 - h) reacting the allylic chloride with diphenylphosphide and then with hydrogen peroxide to form a phosphine oxide;
 - i) reacting the phosphine oxide with phenyllithium to produce a conjugate base of the phosphine oxide;
 - j) coupling the conjugate base of phosphine oxide with a C,D-ring ketone to produce O-silylated derivatives of diastereomeric 2-(4'-hydroxylated) calcitriol analogues; and
- 25 k) separating and isolating the analogues.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/07595

A. CLASSIFICATION OF SUBJECT MATTER				
IPC(6) :C07C 401/00 US CL :552/653 FROM the street of the				
US CL :552/633 According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIEL	DS SEARCHED			
Minimum d	ocumentation searched (classification system followed	by classification symbols)		
	552/653; C07C 401/00			
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)				
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.	
P, Y	US, A, 5,389,622 (POSNER ET AL entire document.) 14 February 1995, see	1-4	
x	JP, A, 6-41059 (NAKASOTO SEIYAKU KK) 15 February			
- Y	1994, see entire document.		1-6	
P, X	1-4			
Further documents are listed in the continuation of Box C. See patent family annex.				
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